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An oxide magnetic material.

An oxide magnetic material provided by the present invention contains, as main components, 55 to 59 mol% of Fe₂O₃; 35 to 42 mol% of MnO; and 6 mol% or less of ZnO, and further contains, as sub-components, 0.05 to 0.3 wt% of CaO; and 0.005 to 0.05 wt% of SiO₂. The other oxide magnetic material provided by the present invention contains, as main components, 61 to 67 mol% of Fe₂O₃; 3 to 36 mol% of MnO; and 30 mol% or less of ZnO, and further contains, as sub-components, 0.05 to 0.5 wt% of CaO; and 0.005 to 0.2 wt% of SiO₂. Such materials can further contain one or more kinds of oxides selected from the group consisting of ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃. Such magnetic materials of the present invention have the advantages of having an extremely low magnetic loss even when used in the high frequency band and having a minimum magnetic loss at a temperature sufficiently higher than room temperature. Moreover, a switching power supply having a switching frequency of 300 kHz to 5 MHz is provided by using such magnetic materials as a magnetic core.

Fig. 1A

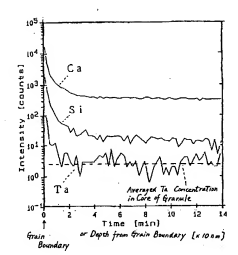
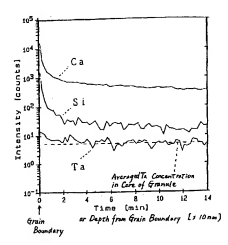


Fig. 1B



BACKGROUND OF THE INVENTION

1. Field of the Invention:

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The present invention relates to a magnetic material. More particularly, it relates to a ferrite type magnetic material having a low magnetic loss when used in a high frequency band.

2. Description of the Related Art:

As electronic technique has been developed recent years, electronic apparatuses have not only become compact and of high density but have become also begun to be used at a higher frequency for improving their efficiencies. For example, a magnetic material used in a magnetic core of a converter in a switching power supply, an inductance component and the like have been required to be compact as well as to be applied for use at a higher frequency. Such a material which has a low magnetic loss when used in a high frequency band has been desired so as to prevent heat generation caused especially in a compact apparatus.

As an example of the prior art, magnetic materials used in a magnetic core and the like will now be described. The magnetic materials are categorized into two major types, metal type materials and oxide ferrite type materials. The metal type materials have advantages of having a high saturation magnetic flux density and a high magnetic permeability. But they have a disadvantage that magnetic loss due to an eddy current is increased when used in the high frequency band since they have a low electric resistivity of about 10^{-6} to 10^{-4} $\Omega \cdot cm$. As this disadvantage can be overcome by making the magnetic material thin, a metal is shaped into a thin foil and rolled up together with an insulator placed thereon. However, the lower limit of the thickness of the metal type materials is about 10 μ m, which is also the lower limit in the abovementioned reduction of an eddy current. Moreover, the metal type materials are disadvantageously difficult to be formed into complicated shapes and expensive. Thus, such materials can not be used in a frequency band over 100 kHz.

Alternatively, the ferrite type materials have a low saturation magnetic flux density of about a half of that of the metal type materials. However, the electric resistivity thereof is much higher than that of the metal type materials: for example, a generally used MnZn type ferrite has an electric resistivity of about 1 $\Omega \cdot$ cm. Moreover, the electric resistivity can be further increased up to about 10 to several hundreds $\Omega \cdot$ cm by adding CaO, SiO₂ and the like. As a result, the magnetic loss due to an eddy current is small and such materials can be used in a relatively high frequency band. In addition, the ferrite type materials have advantages that they can be easily formed into complicated shapes and are inexpensive. Therefore, the ferrite type materials have been generally used in a magnetic core of a converter in a power supply that is used at a switching frequency of, for example, 100 kHz or more and 500 kHz or less.

However, the ferrite type materials can not be used at a frequency of 500 kHz or more because the magnetic loss due to an eddy current is increased at such a high frequency.

When the temperature coefficient of the magnetic loss is positive at around room temperature, the magnetic core emits heat due to the magnetic loss while in use, resulting in a raised temperature. As the temperature rises, the magnetic loss is further increased to make the magnetic core emit more heat. This cycle is repeated, and as a result, a thermorunaway may occur. Therefore, the ferrite type materials need to have a temperature characteristic that the temperature coefficient of the magnetic loss is negative at around room temperature and the magnetic loss is minimized at a temperature where the ferrite type materials are actually used, that is, from room temperature to 80°C. At present, there is no MnZn type ferrite type material that has a sufficiently low magnetic loss when used in a high frequency band. For example, a conventional MnZn type ferrite has a magnetic loss of 1000 kW/m³ or more when used at a frequency of 1 MHz. Moreover, materials which have a comparatively low magnetic loss when used in the high frequency band generally have a bottom temperature of a magnetic loss around room temperature and the temperature coefficient thereof is positive. Thus, a thermorunaway can be easily caused. On the contrary, materials which have a bottom temperature of a magnetic loss over room temperature when used in the high frequency band have a very large magnetic loss. Thus, a material having an extremely low magnetic loss when used in the high frequency band and having a bottom temperature of a magnetic loss sufficiently higher than room temperature has not been provided at the present time. The temperature at which a material has a minimum magnetic loss is referred to as the "bottom temperature of a magnetic loss"

Among magnetic characteristics of a ferrite, characteristics such as the saturation magnetic flux density, the Curie temperature and the bottom temperature of the magnetic loss generally depend upon the

composition of main components. Characteristics such as the magnetic permeability, a residual magnetic flux density, a coercive force and the magnetic loss depend also upon the composition of main components but mainly depend upon the fine structure of the ferrite. Any MnZn type ferrite having a low magnetic loss when used in the high frequency band is required to be high in saturation magnetic flux density, the Curie temperature, the bottom temperature of a magnetic loss and the magnetic permeability (K. Okutani, J. Jpn. Soc. Powder and Powder Metallurgy, 34, (5), p. 191 (1987)).

For example, the saturation magnetic flux density of the MnZn type ferrite is increased when a specific amount of ZnO and a Large amount of Fe₂O₃ are contained therein. However, when the amount of ZnO is too large, the Curie temperature is disadvantageously decreased. In addition, it is known that the magnetic permeability and the electric resistivity is decreased when the amount of Fe₂O₃ is too large. The electric resistivity is decreased because the Fe₂O₃ exceeding 50 mol% is changed into FeO, thereby causing an electron hopping between Fe²⁺ and Fe³⁺. Therefore, as the amount of Fe₂O₃ exceeds 50 mol%, the electric resistivity is decreased. This decrease in the electric resistivity is lowered to some extent by using an appropriate additive or the like. But the thus decreased electric resistivity is still large in comparison with that of a conductive material with a main component originally having a high electric resistivity and comprising an appropriate additive. This decrease of the electric resistivity causes an increase of the magnetic loss due to an eddy current. Therefore, it has been believed that a composition including a large amount of Fe₂O₃ can not be used in the high frequency band.

For example, Japanese Laid-Open Patent Publication No. 61-101458 discloses a MnZn type ferrite comprising 52 to 58 mol% of Fe $_2$ O $_3$, 7 mol% or less of ZnO and 35 to 48 mol% of MnO as main components, and 0.01 to 0.2 wt% of CaO and optionally 0.1 wt% or less of SiO $_2$ as sub-components. This ferrite can be used at a frequency of 100 kHz.

The bottom temperature of a magnetic loss has been considered to depend upon a temperature characteristic of the magnetic permeability. When the MnZn type ferrite is measured for its magnetic permeability against a certain temperature, two peaks are generally obtained. One is a peak at the Curie temperature, which is called a primary peak due to the Hopkinson effect. The other is a peak at around room temperature, which is called a secondary peak. At the temperature of this secondary peak, the crystallomagnetic anisotropic coefficient K_1 is 0. It has been believed that the magnetic loss is a minimal at this temperature.

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The crystallomagnetic anisotropic coefficient K_1 is linearly increased as the temperature rises. When K_1 is negative at room temperature, K_1 is 0 over room temperature. When a composition having such a coefficient is used, the bottom temperature of a magnetic loss can be set over room temperature. K. Ohta, J. Phys. Soc. Japan, 18, p. 684 (1963) describes a change of K_1 obtained by varying a composition ratio of the main components of the MnZn type ferrite. K_1 is slightly affected by the Mn/Zn ratio. K_1 is positive and maximum at room temperature when the ferrite contains about 60 mol% of Fe_2O_3 . K_1 is decreased when the ferrite contains less or more Fe_2O_3 . When the ferrite contains about 55 mol% or less of Fe_2O_3 , or about 61 mol% or more, K_1 is less than 0 at room temperature. Accordingly, the bottom temperature of the magnetic loss can be controlled to be over room temperature when the composition of the ferrite is within the above two ranges. However, when the ferrite contains too much Fe_2O_3 , the magnetic permeability and the electric resistivity are decreased as mentioned above.

Because of the above-mentioned reasons, it has been believed that the most appropriate composition of the main components of the MnZn type ferrite having a low magnetic loss and a high bottom temperature of a magnetic loss is about 53 to 54 mol% of Fe₂O₃, about 9 to 12 mol% of ZnO and the rest of MnO (S. Okamoto, et al., Electronic Ceramics, 16, p. 44 (winter, 1985)). Therefore, most of the ferrites with a low magnetic loss, which have actually been developed up to the present, have the above-mentioned composition. As an approach to decrease the magnetic loss, a method using compositions within or around the above-mentioned range for modifying the kind of additives or the fine structure of the ferrite has been mainly studied. The best ferrite obtained from such an approach has a low magnetic loss of about 500 kW/m³ when used at a frequency of 1 MHz.

For example, Japanese Laid-Open Patent Publication No. 1-224265 discloses a MnZn type ferrite comprising 52.2 to 55.4 mol% of Fe $_2$ O $_3$, 4 to 13.5 mol% of ZnO and 31.1 to 43.8 mol% of MnO as main components, and further comprises at least one sub-component selected from 0.05 to 0.2 wt% of CaO, 0.015 to 0.027 wt% of SiO $_2$, 0.05 to 0.6 wt% of TiO $_2$ and 0.01 to 0.2 wt% of Ta $_2$ O $_5$. This ferrite is sintered from powders having a particle size of 5 μ m or less, and shows a magnetic loss of 300 to 400 kW/m 3 at 1 MHz $_2$ 50 mT.

SUMMARY OF THE INVENTION

The oxide magnetic material which is a sintered substance of this invention comprises, as main components, 55 to 59 mol% of Fe $_2$ O $_3$; 35 to 42 mol% of MnO; and 6 mol% or less of ZnO, and the material further comprises as sub-components, 0.05 to 0.3 wt% of CaO; and 0.005 to 0.05 wt% of SiO $_2$.

In another aspect of the present invention, the oxide magnetic material comprises, as main components, 61 to 67 mol% of Fe₂O₃; 3 to 36 mol% of MnO; and 30 mol% or less of ZnO, and the material further comprises, as sub-components, 0.05 to 0.5 wt% of CaO; and 0.005 to 0.2 wt% of SiO₂.

In still another aspect of the present invention, a switching power supply with a switching frequency of 300 kHz to 5 MHz using this oxide magnetic material as a magnetic core is provided.

Thus, the invention described herein makes possible the advantages of (1) providing a magnetic material having an extremely low magnetic loss even when used in the high frequency band and having a bottom temperature of a magnetic loss sufficiently higher than room temperature, and (2) providing a switching power supply using the magnetic material as a magnetic core and having a switching frequency of 300 kHz to 5 MHz.

These and other advantages of the present invention will become apparent to those skilled in the art upon reading and understanding the following detailed description with reference to the accompanying figures.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A and 1B are profiles of a metal concentration in the direction of depth from a grain boundary of magnetic materials according to the present invention.

Figure 2 is an example of a circuit for a switching power supply using the magnetic material of the present invention.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The inventors of the present invention actually produced various kinds of the MnZn type ferrites by changing composition ratios of main components in a wide range under conditions of adding the same additives without being restricted by the above-mentioned conventional range and studied effects of each composition ratio of the main components. As a result, it was found that a composition including excessive Fe₂O₃ as a main component provides ferrite having a bottom temperature of a magnetic loss of 40 °C or more and having a low magnetic loss when used in the high frequency band of 300 kHz to several MHz, the magnetic loss could be further reduced by using particular additives and adjusting the density of the ferrite.

A sintered substance as a first magnetic material according to the present invention includes 55 to 59 mol% of Fe $_2$ O $_3$, 35 to 42 mol% of MnO and 6 mol% or less of ZnO as main components. The sintered substance further includes 0.05 to 0.3 wt% of CaO and 0.005 to 0.05 wt% of SiO $_2$ as essential subcomponents, and still further contains 0.01 to 0.2 wt% of at least one kind of a third metal oxide M_x O $_z$. The third metal oxide M_x O $_z$ is selected from the group consisting of ZrO $_2$, HfO $_2$, Ta $_2$ O $_5$, Cr $_2$ O $_3$, MoO $_3$, WO $_3$, Al $_2$ O $_3$, Ga $_2$ O $_3$, In $_2$ O $_3$, GeO $_2$, SnO $_2$, Sb $_2$ O $_3$ and Bi $_2$ O $_3$. A sintered substance as a second magnetic material according to the present invention includes 61 to 67 mol% of Fe $_2$ O $_3$, 3 to 36 mol% of MnO and 30 mol% or less of ZnO as main components. This sintered substance further includes 0.05 to 0.5 wt% of CaO and 0.005 to 0.2 wt% of SiO $_2$ as essential sub-components. Preferably, this second magnetic material includes 62 to 66 mol% of Fe $_2$ O $_3$, 14 to 28 mol% of MnO and 10 to 20 mol% of ZnO as main components. Preferably, this sintered substance still further includes 0.01 to 0.5 wt% of at least one kind of a metal oxide M_x O $_z$, which is selected from the group consisting of ZrO $_2$, HfO $_2$, Ta $_2$ O $_5$, Cr $_2$ O $_3$, MoO $_3$, WO $_3$, Al $_2$ O $_3$, Ga $_2$ O $_3$, In $_2$ O $_3$, GeO $_2$, SnO $_2$, Sb $_2$ O $_3$ and Bi $_2$ O $_3$. These first and second magnetic materials were obtained through experiments and study by the inventors of the present invention described below.

According to the study of the inventors of the present invention, the absolute value of the magnetic loss of the MnZn type ferrite largely depended upon the composition ratio of the main components. As the amount of Fe_2O_3 was increased, the magnetic loss was decreased although the magnetic permeability and the electric resistivity were decreased. The absolute value of the magnetic loss could be further reduced by adding effective sub-components to the main components. In other words, a material with a low magnetic loss could be realized by selecting a particular composition of main components and adding a particular amount of CaO and SiO_2 thereto. Further, a material with a lower magnetic loss could be obtained by adding M_xO_z to the main and sub-components. However, if the composition ratio of the main components was not appropriate, addition of an appropriate amount of CaO, SiO_2 and M_xO_z did not provide a material

with a sufficiently low magnetic loss. Moreover, some compositions of the main components could not make the magnetic loss low even with the additives. Therefore, the composition of the main components, especially the amount of Fe_2O_3 , is critical. A material with the lowest magnetic loss can be realized only when an appropriate amount of sub-components are added under an appropriate condition to main components with an appropriate composition ratio.

Next, the design of a magnetic material having a bottom temperature of a magnetic loss higher than room temperature and having the minimum magnetic loss over a temperature at which the material is actually used will now be described. As mentioned above, in the prior art, the bottom temperature of a magnetic loss has been explained by the crystallomagnetic anisotropic coefficient K_I, which varies depending upon temperature. According to this theory, there are two ranges of the composition of the main components in the MnZn type ferrite which can adjust the bottom temperature of a magnetic loss to be over room temperature.

At first, the inventors of the present invention studied a case where 60 mol% or less of Fe_2O_3 was contained. As a result, it was found that the secondary peak of the magnetic permeability was decreased as the amount of Fe_2O_3 was increased as in the conventional theory. However, in the high frequency band over about 1 MHz, the bottom temperature of a magnetic loss did not always correspond to the temperature at which the secondary peak of the magnetic permeability appeared. Actually, the bottom temperature of a magnetic loss was frequently lower than the theoretical value by several tens of degrees centigrade. Thus the range of the usable amount of Fe_2O_3 was narrowed. Especially when 9 to 12 mol% of ZnO was contained, which was conventionally considered to be a suitable composition, the temperature was largely shifted. In this composition, the bottom temperature of a magnetic loss was below room temperature at a frequency of 1 MHz unless the amount of Fe_2O_3 was about 54 mol% or less.

The inventors of the present invention then studied a composition ratio of the main components of the MnZn type ferrite, considering the above described fact. As a result, it was found that the shift of the bottom temperature of a magnetic loss from the secondary peak of the magnetic permeability, that is, the shift from the theoretical value was largely varied depending upon the Mn/Zn ratio even when the same amount of $F_{e_2}O_3$ was contained. When the amount of ZnO was 6 mol% or less, the difference from the theoretical value was decreased as the amount of ZnO was decreased, and sometimes the bottom temperature of a magnetic loss was shifted to be higher than the theoretical value. In the prior art, the amount of $F_{e_2}O_3$ is restricted to 54 mol% or less so that the bottom temperature of a magnetic loss might not be too low. However, it has been impossible to obtain a material with a sufficiently low magnetic loss. In the present invention, it is possible to drastically decrease the magnetic loss and maintain the bottom temperature of magnetic loss at 40 °C or more even when the amount of $F_{e_2}O_3$ is 55 to 59 mol% by decreasing the amount of ZnO from that in the prior art.

Second, the inventors studied a case where 60 mol% or more of Fe_2O_3 was contained. As a result, it was found that K_1 decreased as the amount of Fe_2O_3 was increased, resulting in a raised bottom temperature of magnetic loss. Materials having a composition ratio within this range had lower electric resistivity and lower magnetic permeability as compared with the material containing 53 to 54 mol% of Fe_2O_3 , which is conventionally regarded to be the most appropriate, or a ferrite containing 55 to 59 mol% of Fe_2O_3 . However, the magnetic loss in the high frequency band was very low, which is different from the conventional teachings.

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Further, the inventors studied the decrease of the magnetic loss at the fine structure level by using the additives on the above-mentioned composition of the main components. As a result, it was found that the magnetic loss could be further decreased by adding a small amount of a third metal oxide M_xO_z in addition to the above-mentioned sub-components.

As the third additive, a metal oxide M_xO_z is used. Preferable examples of M_xO_z include ZrO_2 , HfO_2 , Ta_2O_5 , Cr_2O_3 , MoO_3 , WO_3 , Al_2O_3 , Ga_2O_3 , In_2O_3 , GeO_2 , SnO_2 , Sb_2O_3 and Bi_2O_3 . One of the metal oxides or a combination of two or more of them can be used. The added amount thereof is preferably 0.01 to 0.5 wt%.

When a composite additive including four kinds or more of additives, that is, two or more kinds of metal oxides M_xO_z as the third additives in addition to CaO and SiO₂, was used, the effect of minimizing the magnetic loss was also obtained insofar as the total amount of all the used metal oxides exceeded the lower limit of the third additive even if the amount of one of the used metal oxides was less than the lower limit. Of course, when the amount of each metal oxide M_xO_z was within the range, the magnetic loss was decreased as compared with a material in which CaO and SiO₂ alone were used on the main components. Especially, when ZrO_2 and another metal oxide were used together as M_xO_z , not only the magnetic loss was decreased but also the resulting sintered substance was effectively prevented from chipping or breaking. Moreover, addition or containment as impurities of other additives such as TiO₂, CoO, NiO, V₂O₅

and Nb₂O₅ caused no problems insofar as the amount to be added was not too large.

When a large amount of the third additive was used, the electric resistivity was increased and the eddy current loss was decreased. However, when the third additive was dissolved or dispersed in granules of the ferrite and distributed uniformly into the grain boundary and the center of a particle, the hysteresis loss was increased. Therefore, the amount of the third additive to be added must be as small as possible so as to form a thin and uniform deposition in the vicinity of the grain boundary. Thus, a material with a low magnetic loss was obtained. In other words, the magnetic loss was further decreased by making the concentration of the third additive(s) in the grain boundary five times or more as high as the concentration in the inside of the particle. It is known that the eddy current loss can be reduced by adding CaO and SiO₂ at the same time and distributing them in the grain boundary so as to make the electric resistivity higher. The "grain boundary" used herein means the vicinity of the surface of a granule or grain constituting a sintered substance of the ferrite composition. Moreover, it is known that the magnetic loss can be further decreased by using a selected third additive. However, in the prior art, the third additive is not distributed in the grain boundary. According to the study of the fine structure by the inventors of the present invention, it was found to be important to distribute the third additive(s) in the grain boundary.

Further, the inventors found that the density of the sintered substance affected the magnetic loss. Preferably a sintered substance using the MnZn type ferrite with a low magnetic loss has a density of 4.6 g/cm³ or more. If the density of the sintered substance was low, the magnetic loss was increased because an effective cross section was decreased. Moreover, a sintered substance with a low density can be easily affected by the atmosphere when cooled after sintering. In such a case, Fe is reduced. Therefore, especially in a composition including a large amount of Fe₂O₃, it is difficult to obtain essential characteristics of the ferrite without controlling the atmosphere accurately. As a result, the production yield of the sintered substances was decreased. But the yield was improved by making the density 4.6 g/cm³ or more.

The relative magnetic permeability is preferably in a range of 400 or more and 1600 or less. As for the electric resistivity, DC resistance is preferably 100 to 2000 $\Omega \cdot$ cm, and AC resistance (at a frequency of 1 MHz) is preferably about 10 $\Omega \cdot$ cm or more. The magnetic permeability and the electric resistivity vary depending upon grain size. Too small grains reduce the magnetic permeability and too large grains decrease the electric resistivity. An average grain size is preferably 10 μ m or less, and more preferably 2 to 5 μ m.

The MnZn type ferrite material according to the present invention has a bottom temperature of a magnetic loss over 40°C and a low magnetic loss even if measured in a frequency band over 1 MHz. Therefore, a switching power supply with a switching frequency of 300 kHz to 5 MHz using this material as a core of an inverter is compact and of high efficiency and hardly causes any thermorunaway.

Such a magnetic material can be used, as a core of an inverter, in a circuit for a switching power supply, for example, as shown in Figure 2.

Examples

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The present invention will now be described by way of examples.

Example 1

 α -Fe₂O₃, MnCO₃ and ZnO all of which are in the shape of powder and have purity of 99.5% were used as starting materials. Each powder was weighed so as to make a total weight to be 300 g in a composition ratio of each sample as shown in Table 1. All the powder was mixed and wet-ground for 10 hours using a ball mill, and then dried. The mixed powder was calcined in air at 850 °C for 2 hours. CaCO₃ and SiO₂ were added thereto so that the resulting mixture contained 0.1 wt% of CaO and 0.02 wt% of SiO₂. The resulting mixture was again wet-ground for 10 hours by a ball mill, and dried.

Ten wt% of an aqueous solution including 5 wt% of polyvinyl alcohol was added to the calcined powder. The resultant mixture was allowed to pass through a 30 mesh screen to obtain uniform granules. The granules were molded by using a uniaxial molding to give compact bodies. The resulting compact bodies were heated in air at 500 °C for 1 hour to get the binder out, and then sintered under either the following conditions; A or B:

Condition A: The compact body was sintered at a temperature of 1200 °C. The temperature was raised in atmosphere of air. While keeping the highest temperature, atmosphere of oxygen was controlled depending upon an equilibrium oxygen partial pressure for each ferrite. Cooling was performed in a nitrogen rich atmosphere. Concretely, while keeping the highest temperature, a plurality of the compact bodies were respectively sintered at several kinds of oxygen partial pressures in a range of 0.1 to 5% to select the most

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appropriate one. While cooling, a flow rate of nitrogen was gradually increased and the oxygen partial pressure was gradually decreased. The sintering time and pressure during molding were varied so as to make the average grain size of the resulting sintered substance to be about 3 to 5 μ m, the desired density to be about 4.6 to 4.7 g/cm³ and an actual density to be 4.5 to 4.8 g/cm³.

Condition B: The temperature and the atmosphere were the same as in Condition A. The sintering time and pressure during molding were varied so as to make an average grain size of the resulting sintered substance to be 4 μ m or less, the desired density to be about 4.4 g/cm³ and an actual density to be 4.3 to 4.6 g/cm³.

Measurement of a magnetic loss and a bottom temperature of a magnetic loss: A ring-shaped sample having an outer diameter of 20 mm, an inner diameter of 14 mm and a thickness of 3 mm was cut out from each of the resulting sintered substances. Each ring-shaped sample was measured for its magnetic loss under conditions of I MHz•50 mT at 20 °C intervals between 20 °C and 120 °C.

Measurement of a magnetic loss: The ring-shaped ferrite core was coated with one layer of insulating tape. Then the core was wound with an insulated lead wire with a diameter of 0.26 mm. The magnetic loss of the sample was measured by using an alternating current B-H curve tracer. The results are shown in Table 1.

Table 1-1 Effect of main component

	Sample	Composi	tion (mol%)()	Temperature (°	C) ²¹ /Magno	etic Loss (kW/m ³)
5	No.	Fe203	ZnO	MnO	Condition A		Condition B
	1	52	2	46	≥ 120/1420	3)	$\geq 120/1490$
	2	52	3	45	$\geq 120/1290$		$\geq 120/1350$
	3	52	6	42	$\geq 120/1170$		100/1110
10	4	52	10	38	60/ 890		60/ 910
	5	53	2	45	≥ 120/ 960		≥120/ 990
	6	53	3	44	$\geq 120/640$	* 41	100/ 630
	7	53	6	41	100/ 520	*	80/ 500
	8	53	9	38	80/ 630	*	60/ 680
15	9	53	10	37	60/ 710		40/ 720
		54	4	42	100/ 520	*	80/ 550
	10	54	6	40	100/ 420	*	80/ 440
	11	55	0	45	≥ 120/ 880		100/ 920
20	12	55	1	44	100/ 670	*	80/ 900
20	13	, 55	2	43	100/ 440	*	80/ 860
	14	, 55 55	3	42	100/ 290	** ⁵⁾	80/ 550
	15	55	4	41	100/ 280	**	80/ 420
	16	55	6	39	80/ 250	**	60/ 300
25	17	55	8	37	80/460	*	60/ 400
	18	55	9	36	60/ 500	*	60/ 490
	19	55	1.0	35	40/650	*	40/ 620
	20	5 6	4	40	80/ 240	* *	80/ 420
	. 21			38	80/ 230	** .	60/ 480
30	22	56 57	6 0	43	80/ 420	*	80/ 950
	23		1	42	80/ 290	**	60/ 950
	24	57 57	2	41	80/ 200	**	60/ 930
	25		3	40	80/ 180	**	60/ 570
35	26	57	4	39	80/ 190	**	60/ 540
-	27	57 57	5	38	80/ 220	**	60/ 550
	28	57	6	37.	60/ 230	**	60/ 530
	29		7	36	60/ 360	*	60/ 580
	30	57		35	60/ 380		60/ 570
40	31	57	8 9		40/ 430	*	60/ 620
	32	57	_		40/ 490	*	40/ 500
	33	57	10 2		60/ 240	**	≤20/ 990
	34	58	4		60/ 210	**	≤20/ 560
4E	35	58 58	5		60/ 220	**	≦20/ 620
45	36	58	6		60/ 280	**	≦20/ 720
	37	58			≤ 20/ 450		≤20/ 950
	38	58	10	•	40/470	*	≥20/ 300 ≤20/ 980
	39	59	0		40/ 290	**	≦20/ 840 ≦20/ 840
50	40	59	3	38	40/ 430	**	240/ 010

Table 1-2 Effect of main component

Sample	Composi	tion	(mol%) ¹⁾	Temperature (C)2)/Mag	netic Loss (kW/m ³)
No.	Fe203	Zn0	MnÓ	Condition A		Condition B
41	59	4	37	40/ 280	**	≤20/ 710
42	59	6	35	40/ 300	**	≨ 20/800
43	5 9.	9	32	$\leq 20/550$		$\leq 20/1090$
44	60	0	40	$\leq 20/660$	~~~	≤20/930
45	60	3	37	$\leq 20/450$		≤20/960
46	60	6	34	≤20/570		$\leq 20/930$
47	60	15	25	40/ 800		
48	60	30	10	80/ 920		
49	61	0	39	40/1100		
50	61	3	36	60/ 650	*	
51	61	15	24	60/ 540	*	
52	61	30	9	100/ 670	•	
53	61	. 31	8	100/ 880		
54	62	10	28	60/ 250	**	
55	62	20	18	80/ 220	**	
56	64	0	36	60/ 680	*	·
57	64	8	28	60/ 450	*	
58	64	15	21	80/ 150	**	
5 9	64	22	14	100/ 420	*	
		10	24	80/ 250	**	
60	66	20	14	100/ 270	**	
61	66		33	80/ 690	*	
62	67	0	33 18	100/ 520	*	
63	67	15			•	
64	67	30	3	120/ 670	•	
65	67	31	2	120/ 820		
66	68	0	32	80/ 900		
67	88	15	17	100/ 870		
88	68	30	2	120/ 940		

1) In addition to these main components. 0.02 wt% of SiO2 and 0.1 wt% of CaO are contained as sub-components.

2) The bottom temperature of the loss of magnetism (magnetic loss).

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- 4) *: The bottom temperature of magnetic loss is greater than 40° C and magnetic loss is lower than $700 \, (KW/m^3)$.
- 5) **: The bottom temperature of magnetic loss is greater than 40° C and magnetic loss is lower than 300 (KW/m^3).

The results under Condition A shown in Table 1 show that the sintered substances having the following composition of the main components have a bottom temperature of a magnetic loss of 40 °C or more and the magnetic loss was as low as 700 kw/m³ or less: 53 to 59 mol% of Fe₂O₃, 34 to 44 mol% of MnO and 9 mol% or less of ZnO; and 6l to 67 mol% of Fe₂O₃, 3 to 36 mol% of MnO and 30 mol% or less of ZnO. The samples having these excellent properties are marked with (*) in Table 1.

^{3) ---:} The bottom temperature of magnetic loss is lower than 40° C and/or magnetic loss is greater than $700 \text{ (KW/m}^3)$.

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Further, the sintered substances having the following composition of the main components have a bottom temperature of a magnetic loss of $40\,^{\circ}$ C or more and the magnetic loss was as low as 300 kw/m³ or less: 55 to 59 mol% of Fe₂O₃, 35 to 42 mol% of MnO and 1 to 6 mol% of ZnO; and 62 to 66 mol% of Fe₂O₃, 14 to 28 mol% of MnO and 10 to 20 mol% of ZnO. The samples having these excellent properties are marked with (**) in Table 1.

However, under Condition B in which the desired density was about 4.4 g/cm³ and actually was less than 4.6 g/cm³, the resulting sintered substances generally had a higher magnetic loss than the sintered substances obtained under Condition A. The magnetic loss was especially large when 58 mol% or more of Fe₂O₃ was contained therein. The following seems to be the reasons: Because of its low density, (I) the effective cross section was decreased; and (2) the resulting sintered substance was ununiform because the inside of the sintered substance was partially reduced due to the nitrogen rich atmosphere used during cooling. The bottom temperature of a magnetic loss was relatively low in the sintered substances obtained under Condition B as compared with those obtained under Condition A. This seems to be due to the increased amount of Fe⁺² due to the reduction during cooling. Thus, it is preferable that the density of the sintered substance is high.

Example 2

A sintered substance was produced in the composition ratio of each sample shown in Table 2 in the same manner as in Example I under sintering conditions A or B except for the following: $CaCo_3$, SiO_2 and Ta_2O_5 were added so that the resulting mixture contained 0.1 wt% of CaO, 0.02 wt% of SiO and 0.1 wt% of Ta_2O_5 . The resulting sintered substances were measured for the magnetic loss and the bottom temperature of a magnetic loss in the same manner as in Example 1. The results are shown in Table 2.

Table 2-1 Effect of main component

							/+ // 3\
	Sample	Composi	tion	(mol%)1)	Temperature (C)27/Magn	etic Loss (kW/m ³)
5	No.	Fe203	Zn0	MnO	Condition A		Condition B
	1	52	2	46	$\geq 120/1390$	3)	≥ 120/1480
	2	52	3	45	$\geq 120/1220$		$\geq 120/1320$
	3	52	6	42	$\geq 120/1140$		100/1090
10	4	52	10	38	60/ 790		60/850
	5	53	2	45	≥120/ 880	~	≥ 120/ 920
	6	53	3	44	$\geq 120/550$	* 41	100/ 590
	7	53	6	41	100/ 430	*	80/ 440
15	8	53	9	38	80/ 550	*	60/ 580
75	9	53	10	37	60/ 520		40/ 650
	10	54	4	42	100/ 440	*	80/ 520
	11	54	6	40	100/ 350	*	80/ 390
	12	55	0	45	≥ 120/820		100/880
20	13	55	1	44	100/ 590	*	80/ 830
	14	55	2	43	100/ 370	*	80/ 800
	15	55	3	42	100/ 220	** ^{\$)}	80/ 470
	16	55	4	41	100/ 190	**	80/ 350
	17	55	6	39	80/ 180	**	63/ 230
25	18	5.5	8	37	80/ 380	*	60/ 360
	19	55	9	36	60/410	* -	60/410
	20	55	10	35	40/ 580	*	40/ 550
	21	56	4	40	80/ 140	**	80/ 390
30	22	56	6	38	80/150	**	60/ 400
	23	57	0	43	80/420	*	80/ 870
	24	57	1	42	80/ 230	**	60/ 850
	25	57	2	41	80/130	**	60/850
	26	57	3	40	80/ 100	**	60/ 500
35	27	57	4	39	80/ 90	**	80/ 470
	28	57	5	38 -	80/ 110	**	60/ 470
	29	57	8	37	60/120	**	60/460
	30	57	7	36	60/ 250	*	60/ 480
40	31	57	8	35	60/310	*	60/ 490
40	32	57	9	34	40/350	*	60/ 560
	33	57	10	33	40/ 420	*	40/ 440
	34	58	2	40	60/ 150	**	≤20/870
	35	58	4	38	60/ 120	**	≤20/ 480
45	36	58	5		60/ 120	**	≤20/510
	37	58	6	36	60/ 220	**	$\leq 20/640$
	38	58	10	32	≤20/390		≤20/890
	39	59	0	41	40/ 380	*	≨20/900
50	40	59	3	38	40/ 210	**	≨20/790

Table 2-2 Effect of main component

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Sample	Composi	tion	(mol%) ¹⁾	Temperature (°	C) ²¹ /Mag	netic Loss (kW/m
No.	Fe203	Zn0	MnO	Condition A		Condition B
41	59	4	37	40/ 200	**	≤20/630
42	5 9	6	35	40/ 230	**	≤20/720
43	59	9	32	≤20/470		$\leq 20/1030$
44	60	٥	40	$\leq 20/550$		≤20/890
45	60	3	37	$\leq 20/330$		$\leq 20/840$
46	60	6	34	≤20/500		$\leq 20/850$
47	60	15	25	40/.630		
48	60	30	10	80/ 720		
49	61	0	39	40/ 980		
50	61	3	36	60/ 580	*	
51	61	15	24	60/ 450	*	
52	61	30	9	100/ 590	*	
53	61	31	8	100/ 710		
54	62	10	28	60/ 180	**	
5.5	62	20	18	80/ 150	**	
56	64	0	36	60/ 570	*	
57	64	8	28	60/ 250	*	
58	64	15	21	80/ 90	**	
59	64	22	14	80/ 220	*	
60	66	10	24	80/ 200	**	
61	66	20	14	100/ 170	**	
62	- 67	0	33	80/ 590		
63	67	15	18	100/460	*	
64	67	30	3	120/ 560	*	
6.5	67	31	2	120/ 720		
66	6.8	0	32	80/ 820		
67	68	15	17	100/ 730		
6.8	58	30	2	120/ 840		

1) In addition to these main components, 0.02 wt% of SiO₂, 0.1 wt% of CaO and 0.1 wt% of Ta₂O₅ are contained.

2) The bottom temperature of the loss of magnetism (magnetic loss).

3) ---: The bottom temperature of magnetic loss is lower than 40° C and/or magnetic loss is greater than $600 \text{ (KW/m}^3)$.

4) *: The bottom temperature of magnetic loss is greater than 40° C and magnetic loss is lower than 600 (KW/m³).

5) **: The bottom temperature of magnetic loss is greater than 40° C and magnetic loss is lower than 230 (KW/m³).

The results under Condition A shown in Table 2 show, as in Example 1, that the sintered substances having the following composition of the main components have a bottom temperature of a magnetic loss of 40°C or more and the magnetic loss was as low as 600 kW/m³ or less: 53 to 59 mol% of Fe₂O₃, 34 to 44 mol% of MnO and 9 mol% or less of ZnO; and 61 to 67 mol% of Fe₂O₃, 3 to 36 mol% of MnO and 30 mol% or less of ZnO. The samples having these excellent properties are marked with (*) in Table 2.

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Further, the sintered substances having the following composition of the main components have a bottom temperature of a magnetic loss of $40\,^{\circ}$ C or more and the magnetic loss was as low as about 200 kW/m³ or less and 90 kW/m³ at the lowest: 55 to 59 mol% of Fe₂O₃, 35 to 42 mol% of MnO and 1 to 6 mol% of ZnO; and 62 to 66 mol% of Fe₂O₃, 14 to 28 mol% of MnO and 10 to 20 mol% of ZnO. The samples having these excellent properties are marked with (**) in Table 2. Comparing these results with those in Example 1, the magnetic loss is decreased by about 100 kW/m³ in Example 2, in which Ta₂O₅ was further added along with CaO and SiO₂.

On the other hand, in the sintered substances obtained under Condition B having a lower density, the magnetic loss is generally larger and the bottom temperature of a magnetic loss is lower than those of the sintered substances obtained under Condition A as shown in Example 1. Therefore, also in cases where these three sub compositions are used, it is preferable that the density of the sintered substance be high.

Next, the resulting sintered substances were broken to observe their broken-out sections. Intergranular fractures were observed in all of the substances. In other words, the substances were broken at boundaries between particles without breaking the particles. An average grain size was about 4 μ m. Samples 14, 17, 20, 26 and 29 obtained under conditions A and B and samples 50, 52, 56, 62 and 64 obtained under Condition A were measured for distribution of Ta elements on the broken-out sections by secondary ion-mass spectroscopy (SIMS). When several tens of points with a small size of 3 x 3 μ m on one sample were analyzed, the Ta concentration deviated from point to point. Therefore, an average concentration in an area with a size of 50 x 50 μ m was obtained in each sample. A profile of Ta elements in the direction of depth from the broken-out section was obtained. It was found, in all the samples, the larger the distance from the broken-out section (i.e., the grain boundary) was, the lower the Ta concentration became, and when the distance was about several to several tens of nm or more, the Ta concentration remained constant. In all samples, the concentration of the grain boundary was about 10 times or more as high as the constant concentration inside the granule.

Example 3

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Powder of Fe_2O_3 , MnO and ZnO was respectively weighed so as to make a total weight to be 300 g in a composition ratio of 56.5 mol% of Fe_2O_3 , 40 mol% of MnO and 3.5 mol% of ZnO in the same manner as in Example 1. The powder was mixed and wet-ground for 10 hours by a ball mill, and then dried. The mixed powder was calcined in air for 2 hours at $800\,^{\circ}$ C. $CaCO_3$ and SiO_2 were added so that the resulting mixture contained CaO and SiO_2 with the respective amounts shown in Table 3. The resulting powder was again mixed and wet-ground for 10 hours by a ball mill, and dried. Sintered substances (a) were made from this calcined powder under Condition A described in Example 1.

Sintered substances (b) were made in the same manner as above except for a composition ratio of 65 mol% of Fe₂O₃, 17 mol% of MnO, 18 mol% of ZnO, and CaO and SiO₂ of the respective amounts as shown in Table 4.

These sintered substances (a) and (b) were measured for the magnetic loss and the bottom temperature of a magnetic loss in the same manner and under the same condition as in Example 1 (1 MHz, 50 mT). As a result, both sintered substances (a) and (b) showed a bottom temperature of a magnetic loss of 80 °C in any combination of the amounts of CaO and SiO₂. The magnetic loss of the sintered substances (a) and (b) are shown in Tables 3 and 4, respectively.

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Table 3

Effect of sub-components												
SiO ₂ (wt%)		0	0.003	0.005	0.02	0.05	0.06					
CaO (wt%)	0 0.03 0.05 0.2 0.3 0.4	1510 1390 1280 1140 1180 1220	1330 970 650 490 520 600	1190 610 290 280 320 550	990 460 240 220 250 530	850 580 260 250 270 550	770 690 570 520 540 610 KW/m ³					

Main components: Fe₂O₃ 56.5 mol%

ZnO 3.5 mol% MnO 40 mol%;

sintering condition: A.

Table 4

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	Effect of sub-components												
SiO ₂ (wt%)	0	0.003	0.005	0.01	0.1	0.2	0.3						
CaO (wt%)	0	1610	1370	1190	990	850	770	700					
040 ()	0.03	1390	970	610	460	580	690	620					
	0.05	1280	650	310	240	270	340	540					
,	0.1	1110	460	220	180	260	280	410					
,	0.2	1140	490	280	220	250	320	520					
	0.5	1180	520	320	240	270	350	670					
	0.6	1220	600	550	530	550	610	820 KW/m ³					

Main components: Fe₂O₃ 65 mol%

MnO 17 mol% ZnO 18 mol%;

sintering condition: A.

As is evident from Tables 3 and 4, when either CaO or SiO₂ alone was used, the resulting sintered substance has a large magnetic loss. the magnetic loss is decreased by a combination of CaO and SiO₂. Especially in the following ranges, the sintered substances have a low magnetic loss of about 300 kW/m³: in the sintered substances (a), $0.05 \le \text{CaO} \le 0.3$ wt% and $0.005 \le \text{SiO}_2 \le 0.05$ wt%; and in the sintered substances (b), $0.05 \le \text{CaO} \le 0.5$ wt% and $0.005 \le \text{SiO}_2 \le 0.2$ wt%.

Example 4

Sintered substances (c) were made under condition A of Example 1 by using 56.5 mol% of Fe₂O₃, 40 mol% of MnO, 3.5 mol% of ZnO as main components, and adding ZrO₂, CaCo₃ and SiO₂ thereto so that the resulting mixture contained 0.05 wt% of ZrO₂ and CaO and SiO₂ with the respective amounts shown in Table 5. Sintered substances (d) were made under Condition A of Example 1 by using 65 mol% of Fe₂O₃, 17 mol% of MnO, 18 mol% of ZnO as main components, and by adding ZrO₂, CaCo₃ and SiO₂ thereto so that the resulting mixture contained 0.05 wt% of ZrO₂ and CaO and SiO₂ with the amounts shown in Table 6. A ring-shaped sample obtained from each of the resulting sintered substances was measured for the magnetic loss and the bottom temperature of a magnetic loss under the same condition as in Example 3. The results showed all samples of the sintered substances (c) had the minimum magnetic loss at a temperature of 80 ° C. the magnetic loss of the sintered substances (c) and (d) are shown in Tables 5 and 6, respectively.

Table 5

	Effect of sub-components on composition containing 3rd additive												
SiO ₂ (wt%) 0 0.003 0.005 0.02 0.05 0.06													
CaO (wt%)	0 0.03 0.05 0.2 0.3 0.4	1490 1220 1170 1110 1130 1190	1220 880 530 400 450 510	1050 490 200 160 210 480	860 390 130 120 150 470	760 480 170 160 180 490	780 550 500 440 460 520 (KW/m³)						

Main components: Fe₂O₃ 56.5 mol%

MnO 40 mol% ZnO 3.5 mol%;

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3rd subcomponent: ZrO₂ 0.05 wt%;

sintering condition: A.

Table 6

	Effect of sub-components on composition containing 3rd addtive													
SiO ₂ (wt%) 0 0.003 0.005 0.01 0.1 0.2 0.3														
CaO (wt%)	0 0.03 0.05 0.1 0.2 0.5 0.6	1580 1250 1080 950 940 1020 1120	1270 870 570 400 440 480 580	1020 550 280 180 210 280 450	860 320 180 120 140 180 470	780 450 210 150 170 230 490	650 570 300 220 280 320 580	620 590 500 380 480 550 770 (KW/m³)						

Main components: Fe₂O₃ 65 mol%

MnO 17 mol% ZnO 18 mol%;

3rd sub-component: ZrO2 0.05 wt%;

sintering condition: A.

As is evident from a comparison between Tables 3 and 4 of Example 2 and Tables 5 and 6 of this example, the sintered substances including the third additive ZrO_2 have a lower magnetic loss in any combination of the amounts of CaO and SiO_2 .

Especially in the following ranges, the sintered substances have a low magnetic loss if the third additive ZrO_2 is added thereto: in the sintered substances (a), $0.05 \le CaO \le 0.3$ wt% and $0.005 \le SiO_2 \le 0.05$ wt%; and in the sintered substances (b), $0.05 \le CaO \le 0.5$ wt% and $0.005 \le SiO_2 \le 0.2$ wt%.

Example 5

Powder of Fe₂O₃, MnO and ZnO was respectively weighed so as to make a total weight to be 300 g in a composition ratio of 56.5 mol% of Fe₂O₃, 39.5 mol% of MnO and 4 mol% of ZnO in the same manner as in Example 1. The powder was mixed and wet-ground for 10 hours by a ball mill, and then dried. The mixed powder was calcined in air for 2 hours at 800 ° C. CaCO₃ and other metal oxides were added thereto so that the resulting mixture contained 0.1 wt% of CaO, 0.02 wt% of SiO₂, and ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ or Bi₂O₃ with the respective amounts shown in Table 7. Sintered substances containing these metal oxides were made under Condition A of Example 1.

Further, sintered substances including the above-mentioned additives were made in the same manner as above except for a composition ratio of 65.5 mol% of Fe₂O₃, 17.5 mol% of MnO, 17 mol% of ZnO.

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A ring-shaped sample obtained from each of the sintered substances was measured for the magnetic loss and the bottom temperature of a magnetic loss under the same condition as in Example 1. The results showed the minimum magnetic loss in any sample was obtained at 80 $^{\circ}$ C. the magnetic loss of the sintered substances including 56.5 mol% of Fe₂O₃ and including 65.5 mol% of Fe₂O₃ are shown in Tables 7 and 8, respectively.

Table 7

	Effect of 3rd Additive												
(wt%)	0	0.005	0.01	0.03	0.05	0.2	0.3						
ZrO ₂	260	250	190	150	120	150	290						
HfO ₂	260	270	200	190	140	210	340						
Ta ₂ O ₅	260	250	190	160	130	160	320						
Cr ₂ O ₃	260	270	220	200	170	210	380						
MoO ₃	260	270	230	200	190	220	400						
WO ₃	260	260	220	190	180	200	370						
Al ₂ O ₃	260	260	210	200	160	190	390						
Ga ₂ O ₃	260	250	190	150	120	160	350						
ln_2O_3	260	250	180	160	120	180	330						
GeO ₂	260	240	190	150	110	150	340						
SnO ₂	260	270	210	190	160	220	370						
Sb ₂ O ₃	260	250	190	170	140	210	320						
Bi ₂ O ₃	260	280	210	200	180	220	390						

Main components: Fe₂O₃ 56.5 mol%

MnO 39.5 mol%

ZnO 4 mol%;

subcomponents: CaO 0.1 wt%

SiO₂ 0.02 wt%;

sintering condition: A.

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Table 8

		Effe	ct of 3rd	Additive	€	•	
(wt%)	0	0.005	0.01	0.1	0.2	0.5	0.6
ZrO ₂	270	260	180	160	120	160	280
HfO ₂	270	250	210	180	130	200	350
Ta ₂ O ₅	270	260	180	170	120	170	320
Cr ₂ O ₃	270	280	220	210	160	220	390
MoO ₃	270	260	240	160	180	210	420
WO ₃	270	250	230	190	170	200	380
Al ₂ O ₃	270	270	220	200	150	190	380
Ga ₂ O ₃	270	250	180	170	130	170	360
In ₂ O ₃	270	270	190	140	160	190	370
GeO ₂	270	250	180	140	100	160	330
SnO ₂	270	260	220	190	170	230	380
Sb ₂ O ₃	270	260	190	180	190	220	340
Bi ₂ O ₃	270	270	200	190	160	240	380

Main components: Fe₂O₃ 65.5 mol%

MnO 17.5 mol%

ZnO 17 mol%;

subcomponents: CaO 0.1 wt%

SiO₂ 0.02 wt%; sintering condition: A.

As is shown in Table 7, in the sintered substances including 56.5 mol% of Fe_2O_3 , those including ZrO_2 , HfO_2 , Ta_2O_5 , Cr_2O_3 , MoO_3 , WO_3 , Al_2O_3 , Ga_2O_3 , In_2O_3 , GeO_2 , SnO_2 , Sb_2O_3 or Bi_2O_3 each in a range of 0.01 to 0.2 wt% have a lower magnetic loss than those including only CaO and SiO_2 as the subcomponents. The sintered substance having the minimum magnetic loss is one containing 0.05 wt% of GeO_2 , which has a magnetic loss of as low as 110 kW/m³.

As is shown in Table 8, in the sintered substances including 65.5 mol% of Fe_2O_3 , those including the third additives each in a range of 0.01 wt% or more and 0.5 wt% or less have a lower magnetic loss than those including only CaO and SiO_2 as the sub-components. The sintered substance having the minimum magnetic loss is one containing 0.2 wt% of GeO_2 , which has a magnetic loss of as low as 100 kW/m³.

Next, a profile in the direction of depth from a broken-out section of each of the added metal elements was measured by using samples having the minimum magnetic loss (i.e., those including the third additives by 0.05 to 0.2 wt%) in the same manner as in Example 2. The results showed, in all of the samples, the concentration of the added metal elements was highest in the grain boundary, was decreased as the distance from the grain boundary was increased, and remained almost constant when the distance was about several tens of nm or more. In all samples, the concentration in the grain boundary was about 10 to several tens of times as high as that in the inside of a granule.

Example 6

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Powder of Fe $_2$ O $_3$, MnO and ZnO was respectively weighed so as to make a total weight to be 300 g in a composition ratio of 57 mol% of Fe $_2$ O $_3$, 4 mol% of ZnO and 39 mol% of MnO in the same manner as in Example 1. CaCO $_3$, SiO $_2$ and Ta $_2$ O $_5$ were added thereto so that the resulting mixture contained 0.1 wt% of CaO, 0.02 wt% of SiO $_2$ and 0.05 wt% of Ta $_2$ O $_5$. The powder was mixed and wet-ground for 10 hours by a ball mill, and then dried. The mixed powder was calcined in air for 2 hours at 800 to 1200 °C. Then, the resulting powder was again mixed and wet-ground for 10 to 20 hours by a ball mill, and dried.

Ten wt% of an aqueous solution including 5 wt% of polyvinyl alcohol was added to the calcined powder. The resultant mixture was allowed to pass through a 30 mesh screen to obtain uniform granules. This powder was molded by using a uniaxial molding to give compact bodies. The resulting compact bodies were heated in air at $1200\,^{\circ}$ C for 5 hours under the same condition as in Example 1 to obtain sintered substances. Samples were obtained in the same manner as in Example 1 except that Ta_2O_5 was added when grinding the powder after the calcination. A ring-shaped sample obtained from each of the

sintered substances was measured for the magnetic loss and the bottom temperature of a magnetic loss in the same manner as in Example 1. As a result, all samples showed the minimum magnetic loss at 80 °C. Moreover, a broken-out section of each sintered substance was observed by an electron microscope to measure the average grain size of the sintered substance. Further, the Ta concentrations in the grain boundary and in the inside of the granule were measured in the same manner as in Example 2 to calculate a ratio of the concentration in the grain boundary to that inside granule. Examples of the sub-component concentration measurement are shown in Figures 1A and 1B. Figure 1A shows the result obtained with a sample having a grain boundary/inside concentration ratio of Ta greater than 5, and Figure 1B shows a result obtained with a sample having the concentration of Ta smaller than 5. The abscissa of the graph represents the depth from the grain boundary and the ordinate represents a metal oxide concentration. The results are shown in Table 9.

Table 9

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	Effect of Ta concentration ratio												
No	Ta addition	Calcination °C	Sinter °C	Molding Kg/cm	Density g/cm ³	Grain Size μm	Magnetic Loss kW/m ³	Ta Conc. Ratio ¹⁾					
1	Non 2)	900	1200	500	4.70	4	220 ⁵⁾						
2	Before 3)	900	1200	500	4.72	4	100 * ⁶⁾	>10					
3	"	1000	1200	500	4.67	4	130 *	9					
4		1100	1200	500	4.62	5	160 *	5					
5	11	1200	1200	500	4.55	5	290	3					
6	n	1200	1200	1500	4.61	5	260	3					
7	After 4)	900	1200	500	4.68	4	110 *	>10					
8	*	1000	1200	500	4.64	4	120 *	>10					
9	**	1100	1200	500	4.60	5	140 *	>10					
10	,,	1200	1200	500	4.53	4	180 *	>10					

- 1) (Ta concentration at grain boundary)/(Ta concentration at inside of granule)
- 2) Control: Ta is not added.
- 3) Ta is added before calcination.
- 4) Ta is added after calcination.
- 5) ---: Magnetic loss is greater than 200 kW/m3.
- 6) *: Magnetic loss is less than 200 kW/m3.

Main components: Fe₂O₃ 57 mol%

MnO 39 mol% ZnO 4 mol%;

subcomponents: CaO 0.1 wt%

SiO₂ 0.02 wt% Ta₂O₅ 0.05 wt%;

sintering condition: A.

As is evident from Table 9, addition of Ta_2O_5 decreases the magnetic loss. But the effect is largely degraded when the ratio of the Ta concentration in the grain boundary to that in the inside of a grain is 5 or less. Moreover, a sintered substance with a density of less than 4.6 g/cm³ (Sample 10 in Table 9) has a slightly larger magnetic loss as compared with other samples 7 through 9, although the ratio of the concentration is over 5. Moreover, the same effect is obtained regardless of whether Ta was added before or after the calcination.

Example 7

Samples were produced by adding Ta before or after calcination in the same manner and in the same composition as in Example 6 except that the composition of the main components were 65 mol% of Fe₂O₃, 22 mol% of MnO and 13 mol% of ZnO. A ring-shaped sample obtained from each of the resulting sintered substances was measured for the magnetic loss and the bottom temperature of a magnetic loss in the same manner as in Example I. As a result, all samples showed the minimum magnetic loss at 80 °C. An average grain size of the sintered substance was measured by observing the broken-out sections of the

sintered substances with an electron microscope. Moreover, the Ta concentrations in the grain boundary and in the inside of the grain were measured in the same manner as in Example 2 to calculate a ratio of the concentration in the grain boundary to that in the inside of the grain. The results are shown in Table 10.

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Table 10

			Effect of	of Ta cond	centration ratio			
No	Ta addition	Calcination °C	Sinter °C	Molding Kg/cm ²	Density g/cm ³	Grain Size μm	Magnetic Loss kW/m ³	Ta Conc. Ratio 1)
1	Non 2)	900	1200	500	4.65	4	250 ⁵⁾	
2	Before 3)	900	1200	500	4.70	4	120 * ⁶⁾	>10
3	11	1000	1200	500	4.68	4	150 *	9
4	#	1100	1200	500	4.65	5	170 *	5
5	. 17	1200	1200	500	4.57	5	310	3
6	"	1200	1200	1500	4.62	5	320	3
7	After 4)	900	1200	500	4.65	4	150 *	>10
8	17	1000	1200	500	4.61	4	130 *	>10
9	17	1100	1200	500	4.65	5	140 *	>10
10	"	1200	1200	500	4.45	4	200 *	>10

- 1) (Ta concentration at grain boundary)/(Ta concentration at inside of granule)
- 2) Control: Ta is not added.
- 3) Ta is added before calcination.
- 4) Ta is added after calcination.
- 5) ---: Magnetic loss is greater than 200 kW/m3.
- 6) *: Magnetic loss is less than 200 kW/m3.

Main components: Fe₂O₃ 65 mol%

MnO 22 mol%

ZnO 13 mol%;

subcomponents: CaO 0.1 wt%

SiO₂ 0.02 wt% Ta₂O₅ 0.05 wt%; sintering condition: A.

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As is evident from Table 10, addition of Ta_2O_5 decreases the magnetic loss. But the effect is largely degraded when the ratio of the Ta concentration in the grain boundary to that in the inside of the grain is 5 or less. Moreover, a sintered substance with a density of less than 4.6 g/cm³ (Sample 10 in Table 10) has a slightly larger magnetic loss as compared with other samples 7 through 9 although the ratio of the concentration is over 5. Moreover, the same effect is obtained regardless whether Ta was added before or after calcination.

Example 8

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A sintered substance (e) was made under Condition A of Example 1 by using mixed powder in a composition of 57 mol% of Fe $_2$ O $_3$, 39 mol% of MnO, 4 mol% of ZnO, 0.1 wt% of DaO, 0.02 wt% of SiO $_2$ and 0.05 wt% of GeO $_2$.

A sintered substance (f) was made under Condition B of Example 1 by using mixed powder in a composition of 55 mol% of Fe $_2$ O $_3$, 39 mol% of MnO, 6 mol% of ZnO, 0.1 wt% of CaO, 0.02 wt% of SiO $_2$ and 0.05 wt% of GeO $_2$.

A sintered substance (g) was made under Condition A of Example 1 by using mixed powder in a composition of 64 mol% of Fe $_2$ O $_3$, 21 mol% of MnO, 15 mol% of ZnO, 0.1 wt% of CaO, 0.02 wt% of SiO $_2$ and 0.05 wt% of GeO $_2$.

A sintered substance (h) was made under Condition A of Example 1 by using mixed powder in a composition of 64 mol% of Fe $_2$ O $_3$, 16 mol% of MnO, 21 mol% of ZnO, 0.1 wt% of CaO, 0.02 wt% of SiO $_2$ and 0.05 wt% of GeO $_2$.

A sintered substance (i) was made under Condition B of Example 1 by using mixed powder in a composition of 52 mol% of Fe $_2$ O $_3$, 38 mol% of MnO, 10 mol% of ZnO, 0.1 wt% of CaO, 0.02 wt% of SiO $_2$

and 0.05 wt% of GeO2.

These sintered substances (e) through (I) were measured for the magnetic loss in the same manner and under the same conditions as in Example 1.

The sintered substance (e) had a density of 4.71 g/cm³ and a minimum magnetic loss of 130 kW/m³ at a temperature of 80°C. The sintered substance (f) had a density of 4.39 g/cm³ and a minimum magnetic loss of 210 kW/m³ at a temperature of 60 °C. The sintered substance (g) had a density of 4.63 g/cm³ and a minimum magnetic loss of 100 kW/m³ at a temperature of 80 °C. The sintered substance (h) had a density of 4.59 g/cm³ and a minimum magnetic loss of 230 kW/m³ at a temperature of 100°C. In this manner, the sintered substances (e) through (h) have compositions preferred in the present invention and are materials having a low magnetic loss.

On the contrary, the sintered substance (i), which has a composition out of the preferred range of the present invention, has a density of 4.41 g/cm³, and has a minimum magnetic loss of 850 kW/m³ at a temperature of 60 ° C.

Each of the five sintered substances was measured for its magnetic loss at its bottom temperature of a magnetic loss under the condition that a product of a magnetic flux density B and a frequency f (B•f) was constantly 50 mT · MHz. (Under this condition, power supply inverters with the same output have a fixed core size.) The results are shown in Table 11.

Table 11

Magnetic Loss vs. Frequency

130

130

260

540

210

220

300

790

100

140

280

560

h

6700

1270

480

230

200

320

830

i

2520

1360

910

850

1070

1190

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Magnetic Loss (kW/m3) Magnetic Flux (mT) Frequency (MHz) е g 250 6930 6400 7050 0.2 167 0.3 1030 1290 1070 100 0.5 400 550 380

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As is evident from Table 11, the sintered substances (e) through (i) showed the minimum magnetic loss at around a frequency of 0.5 to 2 MHz. Comparing the sintered substances (e) through (h) with the sintered substance (i), the former, having the composition preferred in the present invention, are effective in a frequency band of 300 kHZ or more. However, at 10 MHz, even these sintered substances show increased magnetic loss.

Next, an E-shaped core was cut out from each of the sintered substances. A forward type switching power supply circuit (shown in Figure 2) was fabricated by using the E-shaped core. A rise of the temperature due to the magnetic loss was evaluated. The temperature rise in a magnetic core against frequency and magnetic flux density was measured under a determined low-load condition. The results are

shown in Table 12.

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Table 12

	Emission of I	Heat vs. fi	equency	•			
Frequency (MHz)	Magnetic Flux (mT)	Increase in Temperature (°C)					
		е	f	g	h	i	
0.1	50	1	1	1	1	8	
0.2	50	1	3	1	2	11	
0.3	50	-	5	-	4	29	
0.5	50	3	8	4	9	35	
1.0	50	5	12	6	13	43	
1.2	50	7	15	8	16	47	
1.5	50	10	20	10	18	-	
2.0	50	13	24	12	25		
2.2	50	15	32	16	33	-	
1.0	10	1	2	1	2	12	
1.0	20	4	5	3	4	30	
1.0	50	7	_	7	,	".	
1.0	100	12	22	13	21	_	

As is evident from Table I2, assuming that an allowable temperature rise due to a magnetic loss in the core of the inverter to be 25 °C, the temperature is largely raised in the power supply using the sintered substance (i), and thus the sintered substance (i) can not be used at a high frequency. On the contrary, in the power supplies using the sintered substances (e) through (h) having the composition preferred in the present invention, the temperature rise is low. Therefore, they can be used at up to 2 MHz at 50 mT. This is because each of the used materials has a low magnetic loss and, simultaneously, a satisfactory bottom temperature of a magnetic loss. It can be estimated from the results shown in Table 11 that they would be able to be used around 5 MHz since such materials are generally used with a lower magnetic flux density at a higher frequency. A switching power supply of 2 MHz or more is not practical at the present time because a loss in the other parts of the circuit are increased at such a frequency, but such a circuit could be realized if the loss in the other elements is prevented.

As described in the above-mentioned examples, the power supply having a switching frequency of 100 kHz to 2 MHz using the ferrite material of the present invention emits slight heat, has a high efficiency and causes almost no thermorunaway. Moreover, its characteristics are remarkable especially at a frequency of 300 kHz or more, and such a power supply can be used at a frequency up to 5 MHz if the loss in the circuit can be avoided.

As described above, according to the composition of the present invention, a magnetic material having an excellent temperature characteristic for the magnetic loss and a low magnetic loss can be provided by adding at least a certain amount of CaO and SiO₂, and preferably adding a certain amount of M_xO_z to a MnZn type ferrite having a particular main composition. The M_xO_z is at least one selected from the group consisting of ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃.

Various other modifications will be apparent to and can be readily made by those skilled in the art without departing from the scope and spirit of this invention. Accordingly, it is not intended that the scope of the claims appended hereto be limited to the description as set forth herein, but rather that the claims be broadly construed.

Claims

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- 1. An oxide magnetic material which is a sintered substance comprising, as main components, 55 to 59 mol% of Fe₂O₃; 35 to 42 mol% of MnO; and 6 mol% or less of ZnO, the material further comprising, as sub-components, 0.05 to 0.3 wt% of CaO; and 0.005 to 0.05 wt% of SiO₂.
- 2. An oxide magnetic material according to claim I further comprising 0.01 to 0.2 wt% of at least one kind of a metallic oxide selected from the group consisting of ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃.

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- 3. An oxide magnetic material which is a sintered substance comprising, as main components, 61 to 67 mol% of Fe₂O₃; 3 to 36 mol% of MnO; and 30 mol% or less of ZnO, the material further comprising, as sub-components, 0.05 to 0.5 wt% of CaO; and 0.005 to 0.2 wt% of SiO₂.
- 4. An oxide magnetic material according to claim 3, wherein the main components are 62 to 66 mol% of Fe₂O₃, 14 to 28 mol% of MnO; and 10 to 20 mol% of ZnO.
 - 5. An oxide magnetic material according to claim 3 further comprising 0.01 to 0.5 wt% of at least one kind of a metallic oxide selected from the group consisting of ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃.

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- 6. An oxide magnetic material according to claim 4 further comprising 0.01 to 0.5 wt% of at least one kind of a metallic oxide selected from the group consisting of ZrO₂, HfO₂, Ta₂O₅, Cr₂O₃, MoO₃, WO₃, Al₂O₃, Ga₂O₃, In₂O₃, GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃.
- 7. An oxide magnetic material according to claim 2, wherein a concentration of the metallic oxide in a grain boundary of a grain constituting the oxide magnetic material is 5 times or more as high as a concentration of the metallic oxide in an inside of the grain.
- 20 8. An oxide magnetic material according to claim 5, wherein a concentration of the metallic oxide in a grain boundary of a grain constituting the oxide magnetic material is 5 times or more as high as a concentration of the metallic oxide in an inside of the grain.
- 9. An oxide magnetic material according to claim 6, wherein a concentration of the metallic oxide in a grain boundary of a grain constituting the oxide magnetic material is 5 times or more as high as a concentration of the metallic oxide in an inside of the grain.
 - 10. An oxide magnetic material according to any of claims 1 through 9, wherein an average grain size of a grain constituting the oxide magnetic material is $10 \, \mu m$ or less.
 - 11. An oxide magnetic material according to any of claims 1 through 9, wherein an average grain size of a grain constituting the oxide magnetic material is 2 to 5 µm.
- 12. An oxide magnetic material according to any of claims 1 through 9, wherein a density of the oxide
 magnetic material is 4.6 g/cm³ or more.
 - 13. A switching power supply with a switching frequency of 300 kHz to 5 MHz using the oxide magnetic material according to any of claims 1 through 9 as a magnetic core.



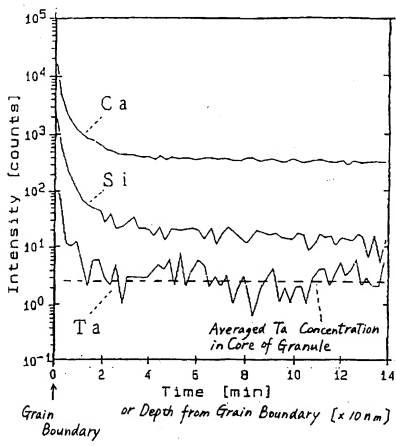


Fig. 1B

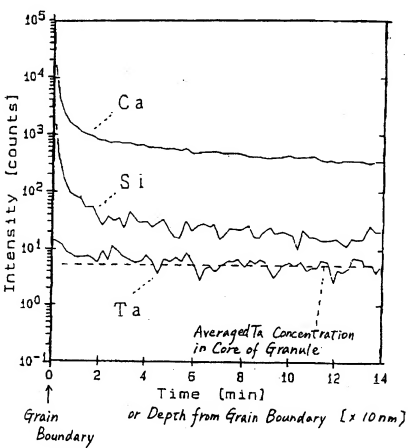
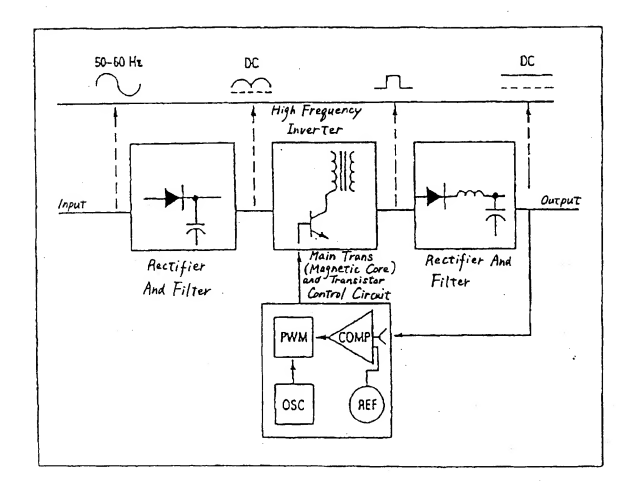


Fig. 2



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(12)

EUROPEAN PATENT APPLICATION

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An oxide magnetic material.

(57) An oxide magnetic material provided by the present invention contains, as main components, 55 to 59 mol% of Fe₂O₃; 35 to 42 mol% of MnO; and 6 mol% or less of ZnO, and further contains, as subcomponents, 0.05 to 0.3 wt% of CaO; and 0.005 to 0.05 wt% of SiO₂. The other oxide magnetic material provided by the present invention contains, as main components, 61 to 67 mol% of Fe₂O₃; 3 to 36 mol% of MnO; and 30 mol% or less of ZnO, and further contains, as sub-components, 0.05 to 0.5 wt% of CaO; and 0.005 to 0.2 wt% of SiO2. Such materials can further contain one or more kinds of oxides selected from the group consisting of ZrO₂, HfO₂, Ta_2O_5 , Cr_2O_3 , MoO_3 , WO_3 , Al_2O_3 , Ga_2O_3 , In_2O_3 , GeO₂, SnO₂, Sb₂O₃ and Bi₂O₃. Such magnetic materials of the present invention have the advantages

of having an extremely low magnetic loss even when used in the high frequency band and having a minimum magnetic loss at a temperature sufficiently higher than room temperature. Moreover, a switching power supply having a switching frequency of 300 kHz to 5 MHz is provided by using such magnetic materials as a magnetic core.

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Fig. 1 A

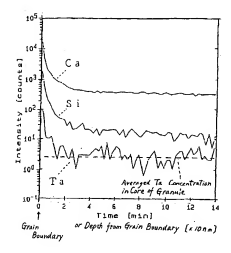
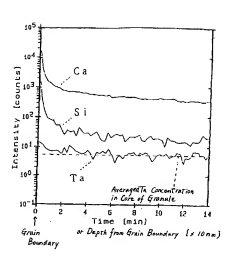


Fig. 1B





EUROPEAN SEARCH REPORT

Application Number

EP 93 10 0497

1		DERED TO BE RELEVA	N 1	
Сатедогу	of relevant pas	dication, where appropriate, sages	Relevant to claim	CLASSIFICATION OF TH APPLICATION (Int.Cl.5)
Х	DE-B-13 02 342 (SIE	1,2,12,	C04B35/26	
	* column 5, line 9 - figure 2 *	- line 13; claims 1,6;		
X.	DE-A-20 22 778 (NIPP LTD.) * claims 1-4 *	PON ELECTRIC COMPANY	1,2,13	
ŀ	GB-A-2 220 935 (MITS		1,2,10, 11,13	
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	The present search report has been	drawn up for all claims		
	face of search	Date of completion of the search		Examiner
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X : particul Y : particul docume A : technol	FEGORY OF CITED DOCUMENTS larly relevant if taken alone larly relevant if combined with another ent of the same category ogical background	E: earlier patent docu after the filing dat D: document cited in L: document cited for	underlying the in- iment, but publish e the application	vantion
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	CLAIMS INCURRING FEES
The pre	sent European patent application comprised by
	sent European patent application comprised at the time of filing more than ten claims.
	All claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for all claims.
	Only part of the claims lees have been paid within the prescribed time limit. The present European search
	report has been drawn up for the first ten claims and for those claims for which claims fees have been paid.
	namely claims:
	No claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims.
TL	ACK OF UNITY OF INVENTION
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]	Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid.
	namely claims:
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د	None of the further search fees has been paid within the fixed time limit. The present European search report
	has been drawn up for those parts of the European patent application which relate to the invention first



EPO FORM 1503 03.82 (POLC01)

EUROPEAN SEARCH REPORT

Application Number EP 93 10 0497

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EUROPEAN SEARCH REPORT

Application Number

EP 93 10 0497

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LACK OF UNITY OF INVENTION A PRIORI

The Search Division considers that the present European petent application does not comply with the requirement of unity of invention and relates to several inventions or groups of inventions.

1. Claims 1,2,7 and 10-13 as far as they are dependent on claims 1,2,7:

Oxide magnetic material with first composition.

2. Claims 3-6.8.9 and 10-13 as far as they are dependent on claims 3-6.8.9:

Oxide magnetic material with second composition.